

Gene expression

Evaluating reproducibility of differential expression discoveries in microarray studies by considering correlated molecular changesMin Zhang^{1,†}, Lin Zhang^{2,†}, Jinfeng Zou¹, Chen Yao², Hui Xiao¹, Qing Liu¹, Jing Wang¹, Dong Wang¹, Chenguang Wang¹ and Zheng Guo^{1,2,*}¹College of Bioinformatics Science and Technology, Harbin Medical University, Harbin 150086, China and²Bioinformatics Centre and School of Life Science, University of Electronic Science and Technology of China, Chengdu, 610054, China

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ABSTRACT

Motivation: According to current consistency metrics such as percentage of overlapping genes (POG), lists of differentially expressed genes (DEGs) detected from different microarray studies for a complex disease are often highly inconsistent. This irreproducibility problem also exists in other high-throughput post-genomic areas such as proteomics and metabolism. A complex disease is often characterized with many coordinated molecular changes, which should be considered when evaluating the reproducibility of discovery lists from different studies.

Results: We proposed metrics percentage of overlapping genes-related (POGR) and normalized POGR (*n*POGR) to evaluate the consistency between two DEG lists for a complex disease, considering correlated molecular changes rather than only counting gene overlaps between the lists. Based on microarray datasets of three diseases, we showed that though the POG scores for DEG lists from different studies for each disease are extremely low, the POGR and *n*POGR scores can be rather high, suggesting that the apparently inconsistent DEG lists may be highly reproducible in the sense that they are actually significantly correlated. Observing different discovery results for a disease by the POGR and *n*POGR scores will obviously reduce the uncertainty of the microarray studies. The proposed metrics could also be applicable in many other high-throughput post-genomic areas.

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1 INTRODUCTION

The reproducibility problem is one of the fundamental issues in validation of high-throughput biological discoveries (Marshall, 2004; Ransohoff, 2004; Ransohoff, 2005) because it is often considered that 'a study that cannot be reliably reproduced has little value' (Klebanov *et al.*, 2007). However, lists of differentially expressed genes (DEGs) detected from different microarray studies for a disease are often highly inconsistent (Ein-Dor *et al.*, 2005;

Michiels *et al.*, 2005; Tan *et al.*, 2003), which frequently raises doubts about the reliability of microarrays (Ein-Dor *et al.*, 2005; Frantz, 2005; Miklos and Maleszka, 2004). It has been suggested that many factors such as large biological variations, small sample sizes (Ein-Dor *et al.*, 2005; Ein-Dor *et al.*, 2006; Zhang *et al.*, 2008) and improper statistical methods (Klebanov *et al.*, 2007; Shi *et al.*, 2006; Tong *et al.*, 2006) may produce inconsistent DEG lists for a disease. Recently, we demonstrated that in the presence of small technical noise and wide differential expressions in a cancer, it is still highly likely that the observed percentage of overlapping genes (POG) of DEG lists from two studies is low, although each DEG list may comprise mostly true DEGs (Zhang *et al.*, 2008).

Up to now, the concept of reproducibility of DEG lists for a disease is often intuitively defined, using terms such as concordance (Shi *et al.*, 2006), agreement (Shi *et al.*, 2006), stability (Ein-Dor *et al.*, 2006; Qiu *et al.*, 2006) or rediscovery rate (Xu and Li, 2003). Most metrics for evaluating the consistency between two gene lists, including the frequently used POG (Ein-Dor *et al.*, 2006; Irizarry *et al.*, 2005; Shi *et al.*, 2006), rely on counting gene overlaps and treat genes as if they would be completely independent with each other. However, a complex disease is often characterized with many coordinated molecular changes (Klebanov *et al.*, 2006; Qiu *et al.*, 2005), which should be taken into account when evaluating whether different studies have actually detected the same or similar results for a disease.

In this article, considering the correlated molecular changes in a complex disease, we propose a novel metric termed percentage of overlapping genes-related (POGR) and its normalized formation *n*POGR for evaluating consistency between two DEG lists for the disease. Obviously, to be utilized as a disease marker, a DEG should have a steady expression pattern in the disease. Thus, we also consider genes' upregulated or downregulated status when evaluating their reproducibility. Based on microarray datasets for three diseases, we show that though there are few genes shared between the DEG lists from different studies for a disease, most of the POGR and *n*POGR scores are rather high, suggesting high reproducibility of the DEG lists in the sense that the apparently inconsistent DEG lists are significantly correlated. After normalizing the effects of list lengths and random data correlations in datasets, the *n*POGR metric may be more appropriate for comparing the

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consistency levels of DEG lists with different lengths and from different studies.

2 METHODS

2.1 Datasets

For prostate cancer, the cDNA microarray data consist of 62 tumors and 41 normal prostate samples (Lapointe *et al.*, 2004), while the oligo microarray (Affymetrix U95Av2) data contain 52 tumor and 50 non-tumor samples (Singh *et al.*, 2002). A total of 6797 genes were measured by both platforms. For lung cancer, the cDNA microarray data consist of 13 squamous cell lung cancer and five normal lung specimens (Garber *et al.*, 2001), while the data by Affymetrix human U95A oligonucleotide arrays consist of 21 squamous cell lung carcinomas and 17 normal lung specimens (Bhattacharjee *et al.*, 2001). A total of 5530 genes were measured by both platforms. The two datasets for Duchenne muscular dystrophy (DMD) are based on Affymetrix (Santa Clara, CA) HG-U95Av2 and HG-U133A GeneChip, respectively. One dataset contains 24 samples from 12 DMD patients and 12 unaffected controls (Haslett *et al.*, 2002) and the other consists of 36 samples from 22 DMD patients and 14 controls (Pescatori *et al.*, 2007). A total of 8572 genes were measured by both platforms. For each disease, we only analyze the genes that are present on both platforms.

The cDNA data is log₂ transformed and then normalized as median 0 and standard deviation 1 per array, as adopted in Oncomine database (Rhodes *et al.*, 2007). The CloneIDs with missing rates above 20% are deleted, and the other missing values are replaced by using the *k*NN imputation algorithm (*k* = 15) (Troyanskaya *et al.*, 2001). The Affymetrix GeneChip data are preprocessed by RMA (Robust Multi-array Analysis). The most recent (July, 2008) SOURCE database (Diehn *et al.*, 2003) is used for annotating CloneIDs to GeneIDs.

2.2 DEGs and co-expression gene pairs

DEGs are most commonly defined as genes with non-zero expression difference between two groups. Usually, each gene is tested against the null hypothesis of equal means of expression levels across groups, while the multiple tests are controlled by false discovery rate (FDR) (Benjamini and Hochberg, 1995) loosely defined as the expected percentage of false positives among the claimed DEGs. When using the conventional *t*-test to select DEGs, small per-gene variances could make genes with small fold-changes statistically significant. To deal with this problem, Tusher *et al.* (2001) proposed the significance analysis of microarrays (SAM) method to select DEGs by adding a small ‘fudge factor’ to the denominator of the *t*-test statistic as follows:

$$d(i) = \frac{\bar{x}_1(i) - \bar{x}_2(i)}{s(i) + s_0} \quad (1)$$

$$s(i) = \sqrt{\frac{1/n_1 + 1/n_2}{n_1 + n_2 - 2} \times \left\{ \sum [x_1(i) - \bar{x}_1(i)]^2 + \sum [x_2(i) - \bar{x}_2(i)]^2 \right\}} \quad (2)$$

where $x_1(i)$ and $x_2(i)$ are the expression vectors of gene *i* in n_1 disease samples and n_2 control samples, respectively. The ‘fudge factor’ s_0 is chosen to minimize the coefficient of variance of $d(i)$.

Because the FDR estimation procedure of SAM might be overly conservative (Xie *et al.*, 2005; Zhang, 2007), we apply the FDR estimation method suggested by (Zhang, 2007) to the SAM program (samr_1.25 R package).

Similarly as Carter *et al.* (2005) did, in each dataset, we estimate the random correlation distribution by independently permuting the sample labels of each gene and then compute the Pearson correlation coefficients for all the gene pairs. For an observed correlation *r* between two genes in the original data, the *P*-value is calculated as the percentage of the permuted correlations exceeding *r*. The FDR control procedure proposed by (Benjamini and Hochberg, 1995) is used to select gene pairs with significantly positive correlation.

2.3 POG and POGR scores

The POG metric is usually used to measure the consistency of gene lists by simply counting overlapped genes between the lists without considering the regulation directions of genes (Chen *et al.*, 2007; Shi *et al.*, 2005). We can evaluate reproducibility of DEG lists in such a broad sense by just counting the overlapped and related genes, as briefly shown in ‘Results’ section (Section 3.3). However, a DEG for a disease should be regulated in the same direction (up-regulated or down-regulated) in different datasets. Thus, strictly, we define a gene is shared by two gene lists only if it is both overlapped and regulated in the same direction in both datasets. In the following text, we mainly analyze the reproducibility of gene lists with such a strict definition.

Suppose *k* genes are shared between list 1 with length l_1 and list 2 with length l_2 , then the POG score from list 1 to list 2 is $POG_{12} = k/l_1$, and the score from list 2 to list 1 is $POG_{21} = k/l_2$.

Then, considering the correlated molecular changes in a disease, we define the POGR metric to evaluate the consistence between two DEG lists. Let O_{r12} (or O_{r21}) represent the number of genes in list 1 (or list 2) which are not shared but significantly positively correlated with at least one gene in list 2 (or list 1), then the POGR scores are calculated as:

$$POGR_{12} = (k + O_{r12})/l_1 \quad \text{or} \quad POGR_{21} = (k + O_{r21})/l_2 \quad (3)$$

Both the POG and POGR scores take values ranging from 0 to 1. Usually, a high POG or POGR score is desired for two discovery lists for a disease. However, both metrics are dependent on the list lengths (see ‘Results’ section) and thus lack statistical significance. Also, when expressions of genes are widely correlated in a disease, the POGR score between two gene lists randomly extracted from the original datasets may be high. To partially solve this problem, for an obtained POGR score, we can estimate the probability (*P*-value) of observing the score in the original datasets by random chance. The null hypothesis is that the POGR score of two DEG lists (with lengths l_1 and l_2) is the same as the expected score for pairs of gene lists (with lengths l_1 and l_2) randomly extracted from the original datasets. Here, the *P*-value is calculated as, among the scores of the 10 000 random pairs of gene lists, the proportion of the scores exceeding the observed one. Similarly to the basic idea of gene set enrichment analysis (GSEA) (Klebanov *et al.*, 2007; Subramanian *et al.*, 2005), which tests whether the members of a gene set are randomly distributed throughout all ranked genes according to correlations of their expressions with the sample labels, we test whether the expression correlations are randomly distributed throughout all genes or primarily among DEGs.

Here, two genes are defined as a correlated pair if and only if they are significantly positively correlated in both datasets. We note that other choices exist. For example, for a DEG list reported in a previous study, it is reasonable to evaluate its reproducibility in a new study by only requiring significant correlation in the new dataset, especially when the original data of the previous study are unavailable. The negatively correlated gene pairs are excluded because we specifically require that the related genes should be regulated in the same direction. However, negatively correlated genes may be also biologically meaningful, which warrants further investigation.

2.4 Normalization of the POG and POGR scores

Because the POG and POGR scores depend on list length, both metrics cannot be applied to compare the consistence levels of pairs of gene lists with different lengths. Thus, we propose an approach to normalize the effect of list lengths on POG and POGR scores. Additionally, we also normalize the POGR scores to reduce the effects of general data correlations in datasets.

For two lists (with lengths l_1 and l_2) randomly extracted from two datasets, suppose $E(k)$ is the expected number of their shared genes, then the POG expectation is $E(POG_{12}) = E(k)/l_1$ or $E(POG_{21}) = E(k)/l_2$. Similarly to the kappa metric (Fleiss, 1971), we normalize the POG score between two DEG lists (with lengths l_1 and l_2) as the proportion of the observed scores beyond

chance to the corresponding maximum potential scores beyond chance as follows:

$$\begin{aligned} nPOG_{12} &= \frac{POG_{12} - E(POG_{12})}{1 - E(POG_{12})} = \frac{k - E(k)}{l_1 - E(k)} \\ nPOG_{21} &= \frac{POG_{21} - E(POG_{21})}{1 - E(POG_{21})} = \frac{k - E(k)}{l_2 - E(k)} \end{aligned} \quad (4)$$

where k is the observed number of the shared genes between the two DEG lists. The $E(POG_{12})$ [or $E(POG_{21})$] can be estimated by the average of the scores for 10 000 pairs of random lists (with lengths l_1 and l_2).

We normalize the POGR in a similar way. For two lists (with lengths l_1 and l_2) randomly extracted from two datasets, suppose $E(O_{r12})$ [or $E(O_{r21})$] is the expected number of genes in list 1 (or list 2), which are not shared but significantly correlated with at least one gene in list 2 (or list 1), then the POGR score expected by random chance is $E(POGR_{12}) = [E(k) + E(O_{r12})]/l_1$ [or $E(POGR_{21}) = [E(k) + E(O_{r21})]/l_2$]. The POGR score can be normalized as:

$$\begin{aligned} nPOGR_{12} &= \frac{POGR_{12} - E(POGR_{12})}{1 - E(POGR_{12})} = \frac{k + O_{r12} - E(k) - E(O_{r12})}{l_1 - E(k) - E(O_{r12})} \\ nPOGR_{21} &= \frac{POGR_{21} - E(POGR_{21})}{1 - E(POGR_{21})} = \frac{k + O_{r21} - E(k) - E(O_{r21})}{l_2 - E(k) - E(O_{r21})} \end{aligned} \quad (5)$$

where $E(O_{r12})$ [or $E(O_{r21})$] can be estimated as the average number of genes in list 1 (or list 2), which are not shared but significantly positively correlated with genes in list 2 (or list 1) in 10 000 pairs of gene lists (with lengths l_1 and l_2) randomly extracted from two datasets.

When the POGR score of two DEG lists (with lengths l_1 and l_2) is less than the expected score for pairs of gene lists (with lengths l_1 and l_2) randomly extracted from the original datasets, the $nPOGR$ score will be negative. In this situation, we set the score as 0. Thus, the $nPOGR$ score achieves 0 if and only if $POGR \leq E(POGR) < 1$. It achieves 1 if and only if $POGR = 1$ and $E(POGR) < 1$. Note that the score is invalid when $E(POGR) = 1$, such as in the extreme situation that all genes are significantly correlated in both datasets for a disease. However, this does not appear in the datasets for the three diseases under this study and would rarely appear in other real applications. Because an $nPOGR$ score is a ratio, it is better to interpret the score together with the maximum potential agreement beyond chance [$1 - E(POGR)$] it can achieve for pairs of lists with the same lengths from the original datasets.

We can estimate the significance level (P -value) for an observed $nPOGR$ score of two DEG lists (with lengths l_1 and l_2) to test the null hypothesis that this score is the same as the expected score for pairs of gene lists (with lengths l_1 and l_2) randomly extracted from the original datasets. This P -value can be calculated from the same set of 10 000 pairs of random gene lists for calculating the P -value of the corresponding POGR score. According to formula (5), for a pair of gene lists randomly extracted from the original datasets, its $nPOGR$ score will be larger than the observed $nPOGR$ score if and only if its POGR score is larger than the observed POGR score, since the $E(POGR)$ is a constant estimated from the 10 000 random gene list pairs. Thus, estimated as the proportion of the random $nPOGR$ scores exceeding the observed $nPOGR$ score, the P -value of the observed $nPOGR$ score is the same as the P -value of the corresponding POGR score.

After excluding the effects of list lengths and general data correlations on chance agreement, the $nPOGR$ score may be more appropriate for comparing the consistency levels of DEG lists with different lengths and from different studies. On the other hand, because the $nPOGR$ score measures the relative agreement beyond chance, it should be necessary to use the POGR scores to reflect the apparent agreement of DEG lists based on the original data correlation.

3 RESULTS

3.1 Reproducibility of studies for three diseases

For prostate cancer, by SAM with 1% FDR control, we detect 1054 and 1343 DEGs in the datasets of oligo and cDNA microarrays,

respectively. The reproducibility of the two studies is measured in two directions. In one direction, for the list of DEGs detected in the oligo microarray data, we measure its reproducibility in the cDNA microarray data as the percentage of these DEGs, which also appear in the cDNA result. The POG_{12} score is as low as 38% and the $nPOG_{12}$ score is only 30%, intuitively suggesting that most of the results from the oligo microarray data are irreproducible in the cDNA microarray result. Then, considering significantly co-expressed gene pairs with 0.1% FDR control, the $POGR_{12}$ score achieves 90%, intuitively suggesting that the DEG list from the oligo microarray data may be highly reproducible in the cDNA microarray data when considering correlated molecular changes. The corresponding $nPOGR$ is 74%, indicating that after subtracting the POGR score by random chance, the observed score achieves a high proportion of the maximum potential agreement beyond chance (0.38). In another direction, for the DEG list detected in the cDNA data, we measure its reproducibility in the oligo microarray data as the percentage of the detected DEGs, which also appear in the oligo microarray result. For the full DEG lists, although the POG_{21} score is only 30% and the $nPOG_{21}$ is as low as 23%, the $POGR_{21}$ score reaches 90% and the $nPOGR$ is as high as 76%, with the maximum potential agreement value as 0.40.

Similar results can be observed for the lists of the top-ranked DEGs from the two studies. For example, although the two lists of the top-10 DEGs from the two datasets share only three genes, the $POGR_{12}$ score is as high as 90%. The corresponding $nPOGR_{12}$ is 89%, reaching a high proportion of the maximum potential value of 0.94. In another direction, both the $POGR_{21}$ and $nPOGR_{21}$ scores are 100%, indicating that the list of the top 10 DEGs from the oligo microarray data is fully reproducible in the cDNA microarray data when considering correlated molecular changes.

Then, we study the reproducibility of two relatively small-scaled experiments for lung cancer. By SAM with 1% FDR control, we detect 2157 DEGs in the oligo microarray data with 38 samples and only 336 DEGs in the cDNA microarray data with 18 samples. Finding significant correlations with 0.1% FDR control, most of the POGR and $nPOGR$ scores are low and only slightly higher than the corresponding POG and $nPOG$ scores (Table 1). Loosening the FDR level to 5% for significant correlations, the DEG lists from the cDNA microarray data are highly reproducible in the larger oligo microarray data, with all the POGR scores and most of the corresponding $nPOGR$ scores reaching above 90% (all with $P < 2 \times 10^{-3}$) (see Supplementary Table S1). In another direction, all the POGR and $nPOGR$ scores for the lists of the top-ranked 10, 50 and 100 DEGs increase to above 70%. However, for the list of all the 2157 DEGs from the larger oligo microarray data, the $POGR_{12}$ and $nPOGR_{12}$ scores measuring its reproducibility in the smaller cDNA microarray data are only 40% and 34%, respectively. This could be explained by the fact that the smaller samples of the cDNA microarray data may have insufficient statistical power in identifying enough DEGs to characterize the disease.

Finally, for a non-cancer disease (DMD), by SAM with 1% FDR control, we detect 805 and 800 DEGs in two datasets, respectively. For the full lists, the POG and $nPOG$ scores are at a rather high level around 50%. Considering significant correlations with 0.1% FDR control level, the POGR and $nPOGR$ scores in both directions increase to above 60%. Notably, for the top-ranked DEG lists, most POGR and $nPOGR$ scores are above 80%, much higher than the corresponding POG and $nPOG$ scores (Table 1). For example, for the

Table 1. Consistence scores between DEG lists for the same disease when using 0.1% FDR control for significant correlations

Datasets	DEGs	POG	nPOG	POGR	nPOGR	Max ^a	DEGs	POG	nPOG	POGR	nPOGR	Max ^a
From 102 samples to 103 samples ^b						From 103 samples to 102 samples ^b						
Prostate cancer	Top 10	0.30	0.30	0.90	0.89	0.94	Top 10	0.30	0.30	1.00	1.00	0.94
	Top 50	0.14	0.14	0.78	0.69	0.70	Top 50	0.14	0.14	0.92	0.89	0.73
	Top100	0.15	0.14	0.80	0.66	0.60	Top100	0.15	0.14	0.94	0.90	0.63
	1054 vs. 1343	0.38	0.30	0.90	0.74	0.38	1343 vs. 1054	0.30	0.23	0.90	0.76	0.40
From 38 samples to 18 samples ^b						From 18 samples to 38 samples ^b						
Lung cancer	Top 10	0.00	0.00	0.00	0.00	1.00	Top 10	0.00	0.00	0.00	0.00	1.00
	Top 50	0.20	0.19	0.20	0.19	0.99	Top 50	0.20	0.19	0.20	0.19	0.99
	Top100	0.31	0.30	0.32	0.31	0.99	Top100	0.31	0.30	0.31	0.30	0.99
	2157 vs. 336	0.13	0.09	0.13	0.09	0.96	336 vs. 2157	0.82	0.75	0.82	0.75	0.72
From 24 samples to 36 samples ^b						From 36 samples to 24 samples ^b						
DMD	Top 10	0.20	0.20	0.70	0.70	1.00	Top 10	0.20	0.20	0.80	0.80	1.00
	Top 50	0.42	0.42	0.92	0.92	0.99	Top 50	0.42	0.42	0.80	0.80	0.99
	Top100	0.54	0.54	0.94	0.94	0.98	Top100	0.54	0.54	0.85	0.85	0.98
	805 vs. 800	0.53	0.50	0.72	0.68	0.89	800 vs. 805	0.53	0.49	0.65	0.61	0.89

^aMax means the maximum potential agreement beyond chance (see 'Methods' section).

^bTwo datasets for each disease are marked by their sample sizes. 'From dataset1 to dataset2' means that the reproducibility of a DEG list detected in dataset1 is evaluated in dataset2.

list of the top 10 DEGs from the dataset with 24 samples, the POG and *n*POG scores for evaluating its reproducibility in the dataset with 36 samples are both 20%, while the corresponding POGR and *n*POGR scores are both 70%. In another direction, the corresponding POGR and *n*POGR scores are both 80%. Thus, the DEG lists from one of the two studies are highly reproducible in another from the view of correlated molecular changes.

Here, all the POG and POGR scores in Table 1 are significantly higher than the average POG and POGR scores of the random list pairs extracted from the corresponding original datasets ($P < 1 \times 10^{-4}$).

3.2 Factors affecting the scores

As shown in Figure 1, for 10 000 pairs of lists with the same length randomly selected from each dataset, both of the average POG and POGR scores increase as the list length increases.

On the other hand, both the corresponding *n*POG and *n*POGR scores fluctuate around zero as the list length increases (Fig. 1). Thus, compared with their non-normalized counterparts, the normalized metrics *n*POG and *n*POGR are more stable and independent of the list lengths and general data correlations in the datasets. For randomly extracted lists, when using less stringent FDR control levels for selecting significant correlations, the *n*POGR scores are still around zero though the POGR scores may increase (Fig. 2).

As shown in Figure 3, in each of the datasets for prostate cancer, the correlation distribution for all gene pairs is similar to the correlation distribution for DEG pairs selected by SAM with 1% FDR control, indicating that the expressions of genes in the prostate cancer data tend to be widely correlated. In such a situation, the *n*POGR scores decrease obviously from the POGR scores (Table 1). In the two datasets for lung cancer, the distributions of the correlations of DEG pairs are distinct from the distributions

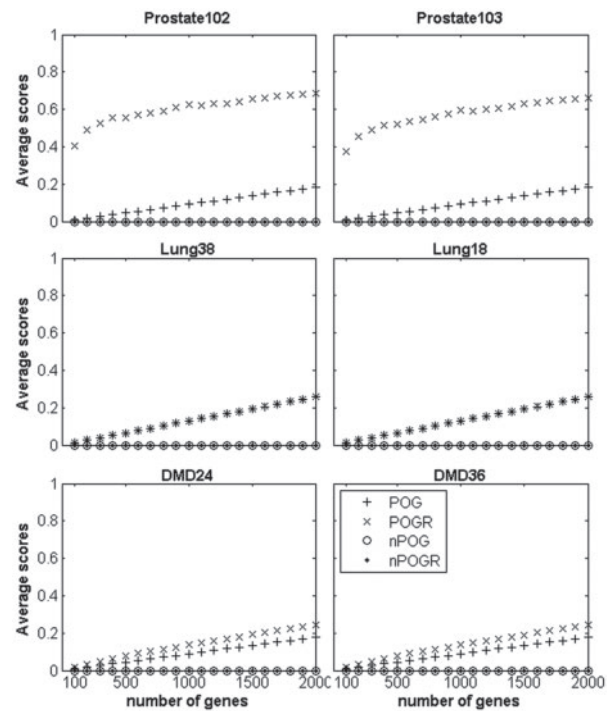


Fig. 1. The effect of list lengths on POG, *n*POG, POGR and *n*POGR. The x-axis represents list lengths ranging from 100 to 2000, and the y-axis represents the average scores of 10 000 pairs of gene lists randomly selected from the original datasets. Here, 0.1% FDR level is used for detecting significant correlations. The legend for all the six subgraphs is the same as shown within the last one.

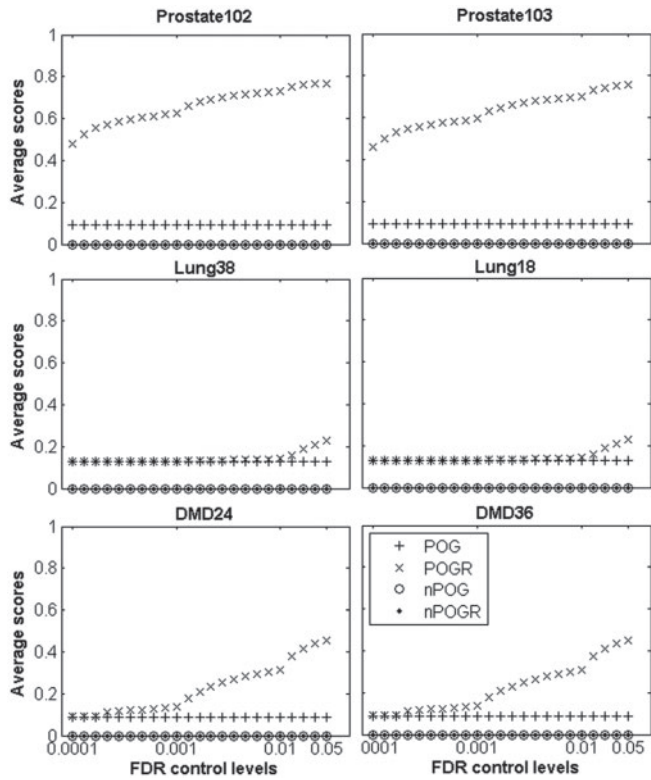


Fig. 2. The effect of FDR control levels on POG, *n*POG, POGR and *n*POGR. The x-axis represents FDR control levels ranging from 0.01% to 5%. The y-axis represents the average scores of 10 000 pairs of gene lists with a given length (1000) randomly selected from the original datasets. The legend for all the six subgraphs is the same as shown within the last one.

of all gene pairs, indicating that the expression correlations in these datasets are largely limited to DEGs. Similar results are observed in the two datasets for DMD. In this situation that most non-DEGs are not significantly correlated, the *n*POGR scores decrease slightly from the POGR scores (Table 1).

3.3 Comparison between DEG lists from different diseases

Finally, we analyze the consistence of DEG lists from datasets for different diseases, using the larger dataset for each disease. The DMD dataset shares a total of 9200 and 8495 genes with the prostate and lung cancer datasets, respectively, and the two cancer datasets have 6797 genes in common. Similarly to the above analysis, DEGs are selected by SAM with 1% FDR control and the correlated gene pairs are determined with 0.1% FDR control.

As shown in Table 2, the POGR and *n*POGR scores of DEG list pairs from different diseases are obviously much smaller than the scores from the same disease. For example, the *n*POGR score is 0 for the full DEG lists from prostate and DMD, while the *n*POGR score is 74% for the full DEG lists from two prostate cancer datasets and 68% for the full DEG lists from two DMD datasets. For DEG lists from a cancer dataset and a DMD dataset, most of the POG and POGR scores are not much different from the scores for gene lists randomly selected from the original datasets ($P > 1\%$). When using looser FDR control levels for selecting correlated genes, the *n*POGR

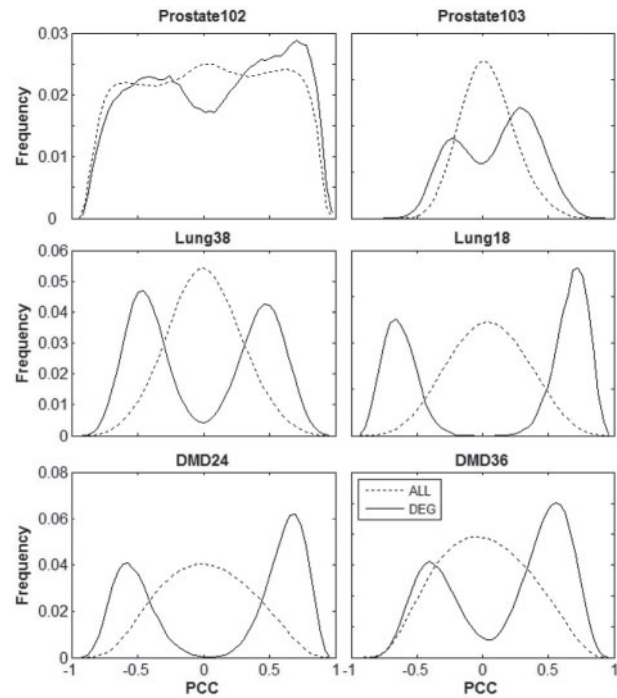


Fig. 3. The distributions of the PCC of expressions of gene pairs in three pairs of datasets. The legend for all the six subgraphs is the same as shown within the last one.

scores of DEG lists from DMD and cancer still fluctuate around zero (Supplementary Table S2).

Notably, the *n*POGR scores between the DEG lists from the two cancers are higher than the scores between DMD and cancer (Table 2 and Supplementary Table S2), suggesting that the expression patterns of the altered genes in different cancers may be more consistent. For example, between the full DEG lists from prostate and lung cancer, the *n*POGR₁₂ score is 15% and the corresponding POGR₁₂ score is 35%, which is significantly higher than the average score of randomly selected lists.

As mentioned in ‘Methods’ section (Section 2.3), we can also evaluate the reproducibility of DEG lists in a broad sense by just counting the overlapped and correlated genes, without regulation direction requirement. Calculated similarly as the approaches described in ‘Methods’ section, for DEG lists selected from the datasets for the same disease, all the scores are almost unchanged (Supplementary Table S3 and Table S4), reflecting that DEGs from the same disease are regulated in a similar way. However, interestingly, for DEG lists from the cancer and DMD datasets, some scores are unexpectedly high (Supplementary Table S5 and Table S6), especially when using less stringent FDR control levels for selecting correlated gene pairs. To understand these surprising results, we find that the full DEG list from DMD are non-randomly overlapped with the full DEG list from the prostate cancer data (POG = 30%, $P < 1 \times 10^{-4}$) as well as the full DEG list from the lung cancer data (POG = 53%, $P < 1 \times 10^{-4}$), indicating that many genes (DEGs) disturbed in DMD are also differentially expressed in cancer. Without the requirement for regulation direction, the overlapped DEGs between DMD and cancer tend to be correlated with other DEGs in each disease and thus will introduce higher POGR as well

Table 2. Consistence scores between DEG lists for different diseases when using 0.1% FDR control for significant correlations

Datasets	DEGs	POG	nPOG	POGR	nPOGR	Max ^a	DEGs	POG	nPOG	POGR	nPOGR	Max ^a
From Prostate to DMD ^b						From DMD to Prostate ^b						
Prostate vs. DMD	Top 10	0.00	0.00	0.00	0.00	0.98	Top 10	0.00	0.00	0.00	0.00	0.98
	Top 50	0.02	0.02	0.04	0.00	0.93	Top 50	0.02	0.02	0.04	0.00	0.93
	Top100	0.01	0.00	0.05	0.00	0.89	Top100	0.01	0.00	0.06	0.00	0.89
	1842 vs. 801	0.05	0.00	0.16	0.00	0.68	801 vs. 1842	0.11	0.01	0.37	0.00	0.52
From Lung to DMD ^b						From DMD to Lung ^b						
Lung vs. DMD	Top 10	0.00	0.00	0.00	0.00	1.00	Top 10	0.00	0.00	0.00	0.00	1.00
	Top 50	0.00	0.00	0.00	0.00	1.00	Top 50	0.00	0.00	0.00	0.00	1.00
	Top100	0.00	0.00	0.00	0.00	0.99	Top100	0.00	0.00	0.00	0.00	0.99
	3303 vs. 797	0.05	0.00	0.05	0.00	0.95	797 vs. 3303	0.21	0.02	0.21	0.02	0.81
From Prostate to Lung ^b						From Lung to Prostate ^b						
Prostate vs. lung	Top 10	0.00	0.00	0.00	0.00	1.00	Top 10	0.00	0.00	0.00	0.00	1.00
	Top 50	0.02	0.02	0.02	0.02	1.00	Top 50	0.02	0.02	0.02	0.02	1.00
	Top100	0.04	0.03	0.04	0.03	0.99	Top100	0.04	0.03	0.04	0.03	0.99
	1353 vs. 2763	0.35	0.15	0.35	0.15	0.77	2763 vs. 1353	0.17	0.06	0.17	0.07	0.89

Indicated as in Table 1.

as *n*POGR scores. However, when considering regulation directions of genes, all the scores become rather low, suggesting that the genes differentially expressed in different diseases may be regulated in different ways. Thus, it is important to consider the regulation directions of DEGs when evaluating their reproducibility.

4 DISCUSSION

Intuitively, a high POG score is expected for two studies for a disease. Similarly, a high POGR score suggests apparently high reproducibility of DEG lists in the sense that different studies have actually detected similar results for a disease. Because both POG and POGR scores are dependent on list lengths, they cannot be simply used to compare the reproducibility of gene lists with different lengths. Statistically, for each obtained POG or POGR score, we can calculate the *P*-value of observing the score in random lists from the original data, which may help interpret the intuitive score.

However, even a small *P*-value for a POG or POGR score cannot guarantee a high percentage of shared genes between two lists. By normalizing the effects of list length and correlation distribution in datasets, the proposed *n*POGR metric suggests a more appropriate way for comparing the reproducibility of gene lists with different lengths and from different studies.

As previously demonstrated by us (Zhang *et al.*, 2008), though DEG lists from current small-scaled microarray studies may only capture a small fraction of DEGs in a disease and thus show low POG scores, each separately determined list may comprise mostly true DEGs. Usually, the global expression changes of genes in a disease may introduce great uncertainty to the findings at the individual genes level. Under such a situation, even if there are many ‘true’ DEGs in a disease, we may be only interested in analyzing the most significant ones. The proposed POGR metric is especially helpful for evaluating such short lists of interesting genes. It should

be noted that distinct methods for selecting DEGs, such as SAM (Tusher *et al.*, 2001), analysis of variance (ANOVA) (Hardeo and Mohammed, 2000) and empirical Bayes with *t*-statistic (Efron *et al.*, 2001; Oshlack *et al.*, 2007), may capture different statistical aspects of expression changes and contribute to the observed inconsistency between the derived DEGs (Jeffery *et al.*, 2006; Qiu *et al.*, 2006). Thus, it warrants further investigation to study the reproducibility of DEG lists found by different DEG selection approaches.

At the functional module level, the strikingly inconsistent gene lists generated in different experiments for a disease could be rather functionally consistent (Guo *et al.*, 2006; Hosack *et al.*, 2003; Zhu *et al.*, 2007). Specifically, we have demonstrated functional modules related to a disease could be captured by only a small fraction of DEGs in the disease (Yang *et al.*, 2008). Therefore, similarly to POGR metric, a metric considering gene overlaps from the view of functional similarity will probably provide another way for evaluating the reproducibility of the apparently inconsistent discoveries in microarray studies, which deserves our future work. Other metrics or views could be considered for observing different discovery results for a disease. Notably, Mao *et al.* (2009) showed that the discriminating genes identified in a study may preserve their disease discriminating ability in another study though these genes might not be selected independently in the later dataset because they might not be the optimum ones in both studies. Their result implies high concordance of DNA microarrays by the signals and classifier transferability of the discriminating genes, rather than simply relying on counting gene overlaps. This concept is somewhat similar to ours. However, our proposed metrics should be more statistically interpretable and applicable in classical studies on evaluating differential gene expressions in diseases.

Finally, we note that the irreproducibility problem of finding disease markers also exists in other post-genomic areas such as proteomics (Ransohoff, 2005), metabolomics (David I. and

Douglas B., 2006) and large-scale screens for cancer mutations (Chen *et al.*, 2007; Frantz, 2005). The apparently inconsistent discoveries generated in these post-genomic areas could also be evaluated by considering correlated disease markers. Because this problem is of fundamental importance in the field of systems biology, it deserves our future researches.

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